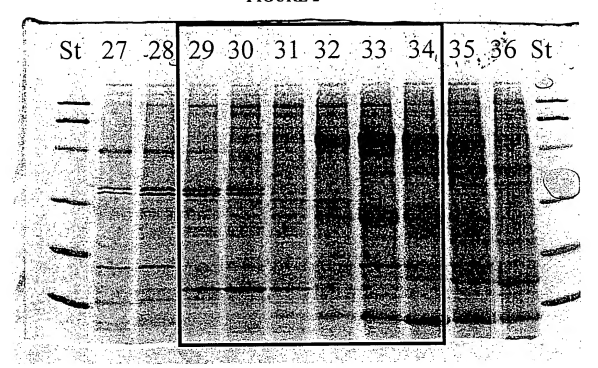
BEST AVAILABLE COPY

SDS – POLYACRYLAMIDE GEL OF OSTEOINDUCTIVELY ACTIVE TO PROTEINS FROM HPLC

Weight		pec
Molecular Weight (kD)	Reduced	Non-reduced
66 —		
45 —		
36 —		
24 —		
	هستندیس آدامیسنیس استنیسی	
20 —		,,
14 —		

FIG. 1



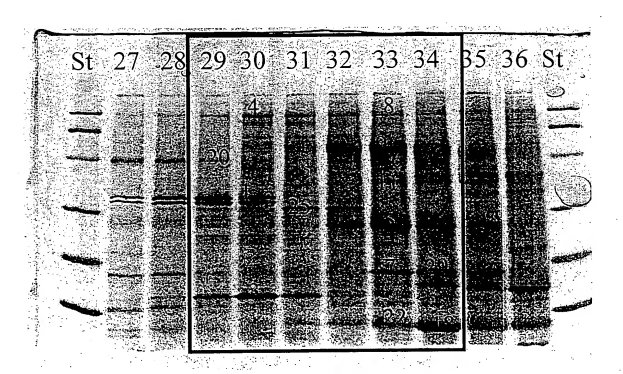
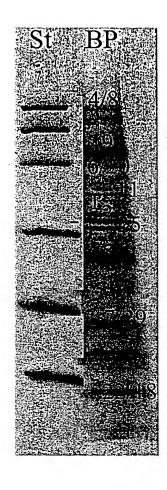


FIGURE 3



Band No.	Identity
1	Histone H1.c
2	Histone H1.c
3	Ribosomal protein RS20
4	Similar to ribosomal protein LORP
5	BMP-3
6	α2 macroglobulin RAP and BMP-3
7	Similar to ribosomal protein LORP
8	BMP-3
9	BMP-3
11	Ribosomal protein RL6 and BMP-3
18	TGF-β2 / SPP 24
20	Factor H
22	TGF-β2
25	BMP-3 and H1.x
29	BMP-3 and ribosomal protein RL32

FIGURE 4

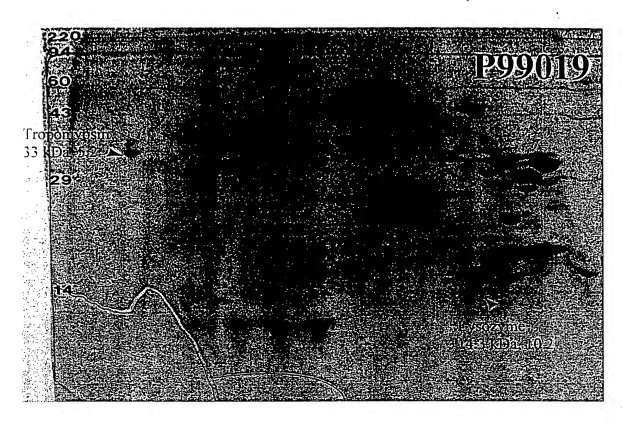


FIGURE 5

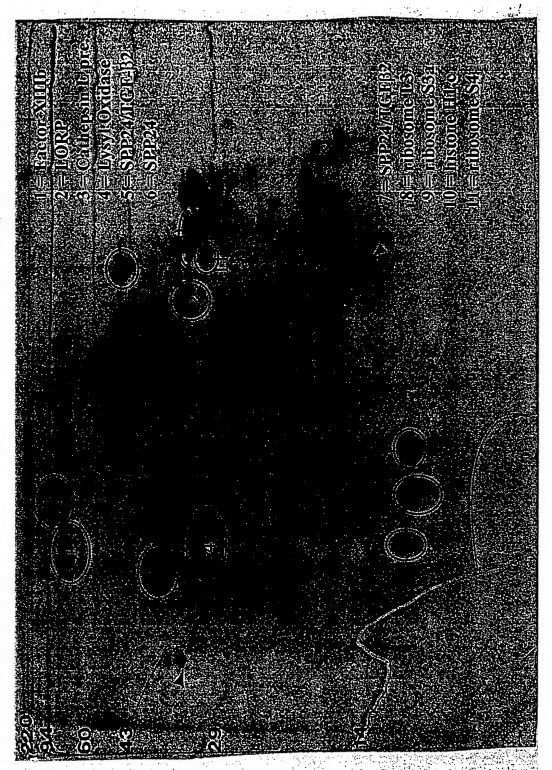
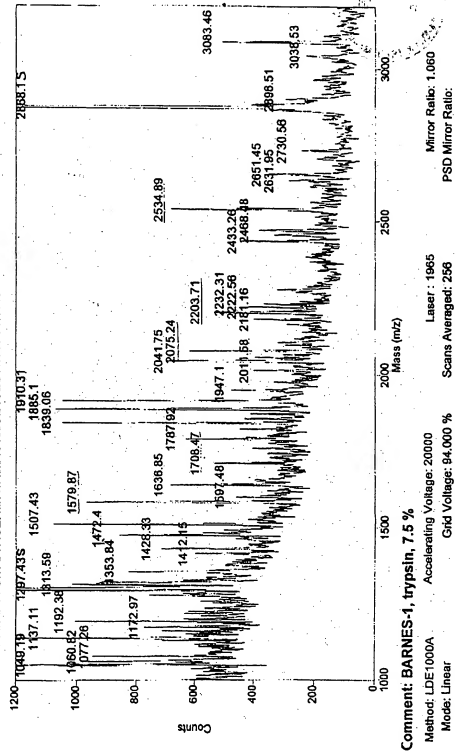


Figure 7A (Band 1)



Original Filename: c:\voyager\data\mag1099\d]gest\barne026.ms
This File # 1 : C:\VOYAGER\DATA\MAG1099\D|GEST\SMOOTH.MS Coll

Savitsky-Golay Order = 2 Points = 19 Collected: 10/12/99 2:13 PM Sample: 74



Timed Ion Selector: 16.1 OFF

Pressure: 1.70e-06

Guide Wire Voltage: 0.075 %

Delay: 50 ON

Low Mass Gate: 500.0

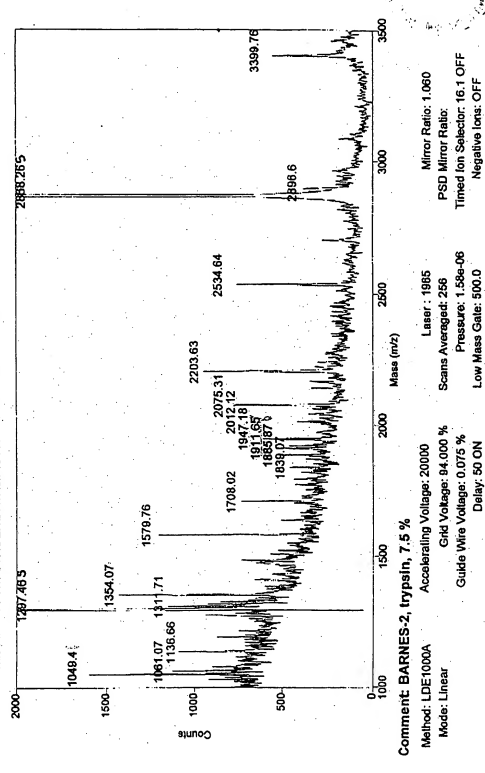
Negative lons: OFF



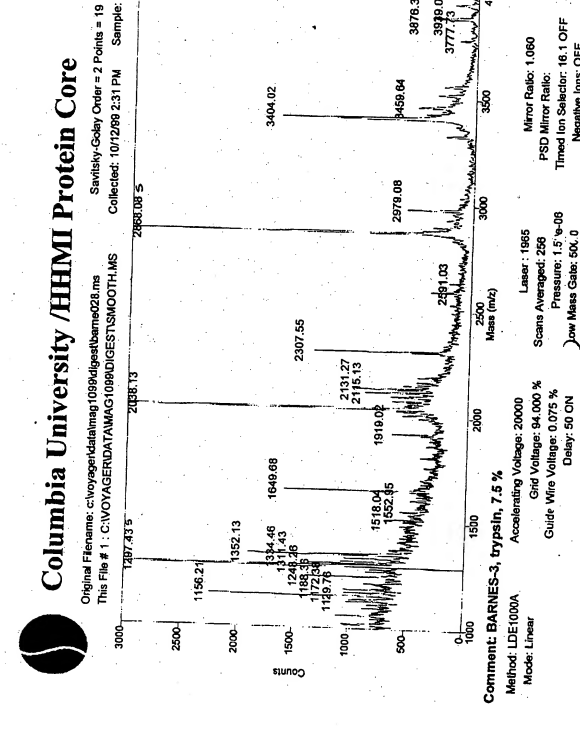
[Protein Core Columbia University /HHMI

Original Filename: c:\voyager\data\mag1099\diges\Dame027.ms This File # 1 : C:\VOYAGER\DATA\MAG1099\DIGEST\SMOOTH.MS

Savitsky-Golay Order = 2 Points = 19 Collected: 10/12/99 2:21 PM Sample: 75



Sample: 76



3876.36

Negative lons: OFF

Figure 7D (Band 4)



Original Filename: c:\voyager\data\mag1099\barnes\barne037.ms This File # 1 : C:\VOYAGER\DATA\WAG1099\BARNES\SMOOTH.MS

Savitsky-Golay Order = 2 Points = 19 Collected: 10/27/99 2:30 PM Sample: 22

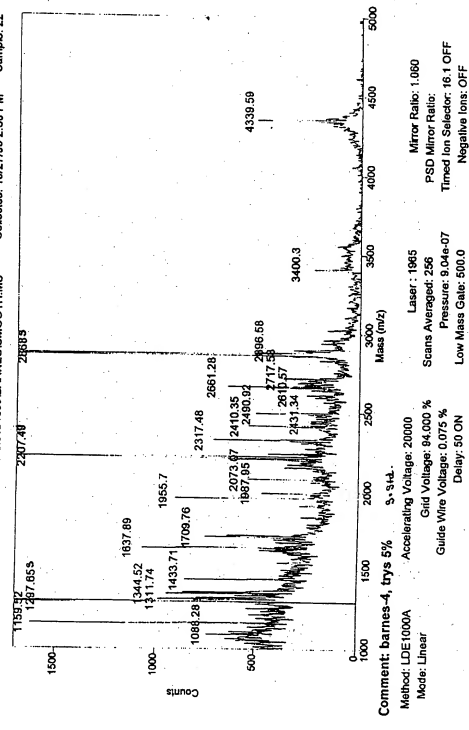


Figure 7E (Band 5)

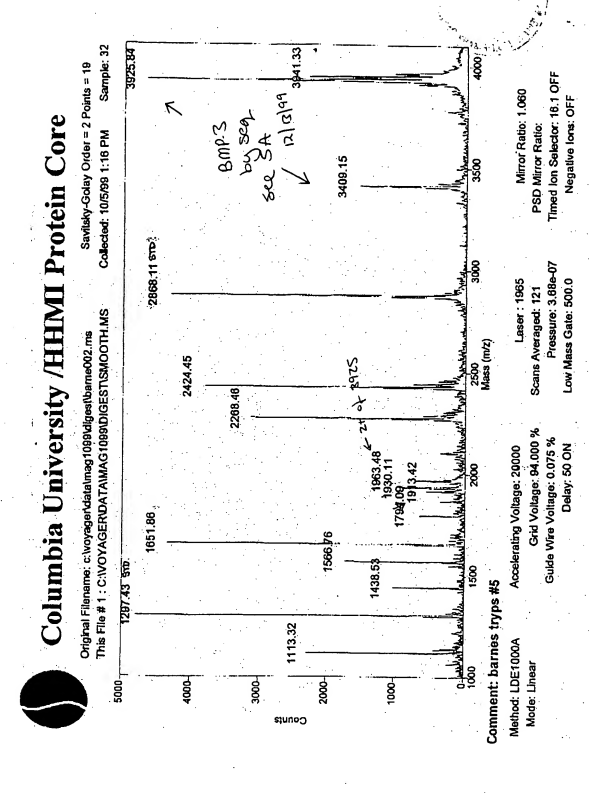


Figure 7F (Band 6)



Original Filename: c:\voyager\data\mag1099\diges\tanson Savits
This File # 1 : C:\VOYAGER\DATA\MAG1099\DIGEST\SMOOTH.MS Collected

Savitsky-Golay Order = 2 Points = 19 Collected: 10/5/99 1:14 PM Sample: 33

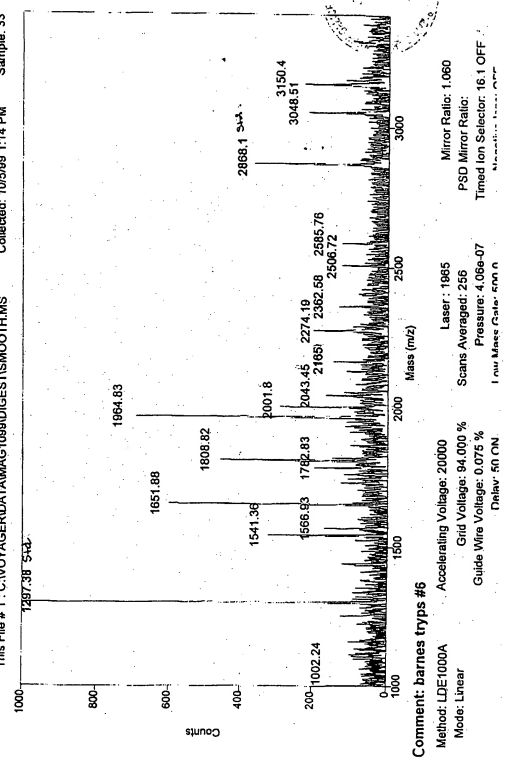
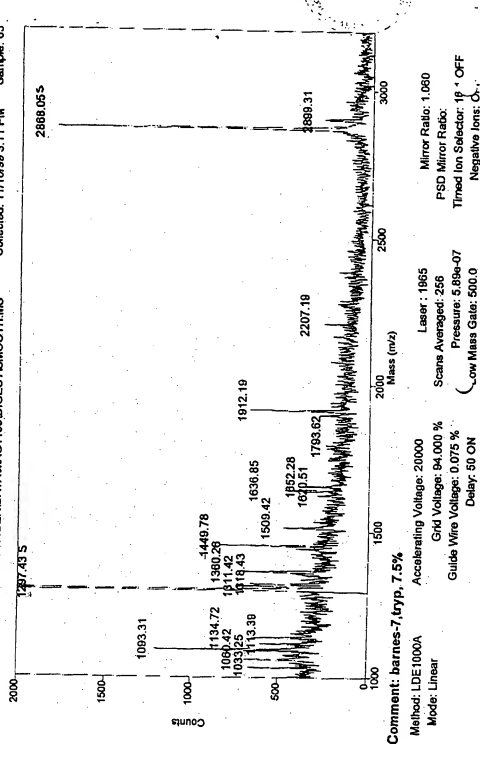


Figure 7G (Band 7)





Sample: 65 Savitsky-Golay Order = 2 Points = 19 Collected: 11/10/99 3:11 PM



Negative Ions: O.

Protein Core Columbia University /HHMI Original Filename: c:\voyager\dala\mag1199\digest\snow_005.ms

Savitsky-Golay Order = 2 Points = 19 Collected: 11/10/99 3:18 PM This File # 1: C: WOYAGERIDATAIMAG11981DHGESTISMOOTH.MS

2000

1500-

1000

Counts

ठु

Sample: 64

2661.45 2410.37 2317.68 2000 Mass (m/z) 2073.11 1955.74 1709.76 1541.59 1159.32

Comment: barnes-8,tryp, 7.5% Method: LDE1000A

.

Accelerating Voltage: 20000

Mode: Linear

Grid Voltage: 94.000 %

Guide Wire Voltage: 0.075 % Delay: 50 ON

Mirror Ratio: 1.060 PSD Mkror Ratio;

Pressure: 4.01e-07

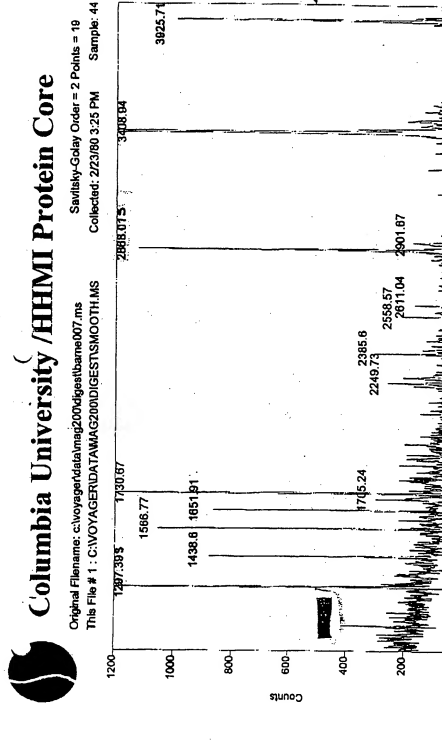
Low Mass Gate: 500.0

Laser: 1965

Scans Averaged: 258

Timed Ion Selector, 16.1 OFF Negative Ions: OFF

Figure 7I (Band 9)



Timed ton Selector. 18.1 OFF Mirror Ratio: 1.080 PSD Mirror Ratio: Pressure: 3.27e-07 Laser: 1860 Low Mass Gate: 500.0 Scans Averaged: 258 Grid Voltage: 94.000 % Guide Wire Voltage: 0.090 % Delay: 50 ON Accelerating Voltage: 25000 Comment: barnes-b9, tryp.mod.,7.5%

2500 Mass (m/z)

2000

1500

96

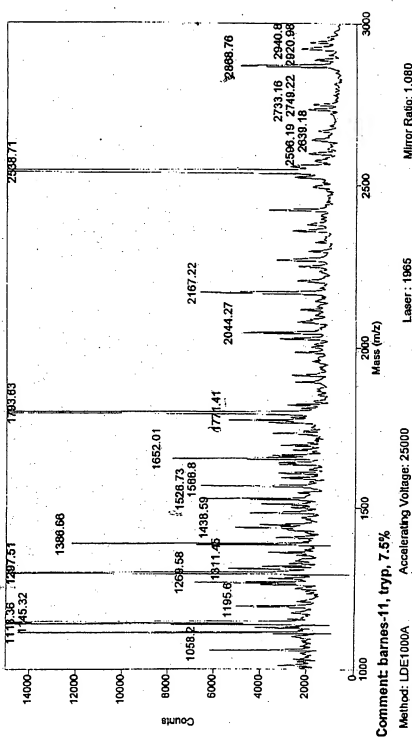
Method: LDE1000A Mode: Linear

Negative Ions: OFF



This File # 2: C:IVOYAGERIDATAIMAG1299DIGESTISMOOTH.MS Original Filename: c:\voyager\data\mag1299\digest\bame004.ms

Savitsky-Golay Order = 2 Points = 19 Collected: 12/15/99 3:49 PM



Accelerating Voltage: 25000

Mode: Linear

Grid Voltage: 94,000 % Guide Wire Voltage: 0.090 %

Delay: 50 ON

Pressure: 5.84e-07 Cow Mass Gate: 500.0 Scans Averaged; 256

Mirror Ratio: 1.080 PSD Mirror Ratio:

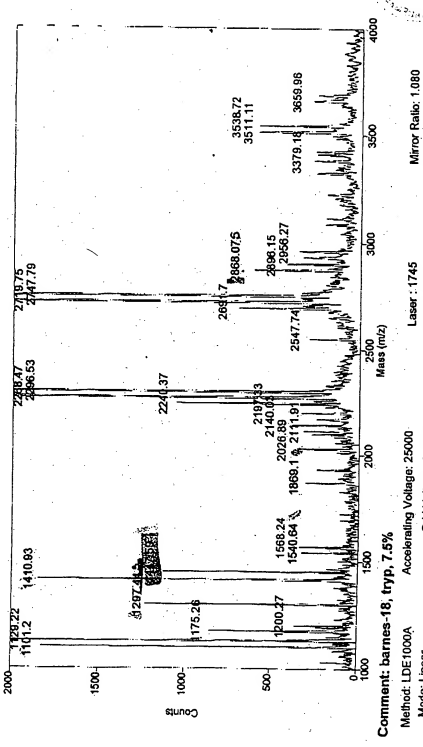
Timed Ion Selector, 19 1 OFF Negative lons: & .



Protein Core Columbia University /HHMI



Savitsky-Golay Order = 2 Points = 19 Collected: 12/15/99 4:47 PM Sample: 13



Timed Ion Selector, 16.1 DFF

Pressure: 2.90e-07

Scans Averaged: 258

Grid Voltage: 94.000 %

Guide Wire Voltage: 0.090 %

Delay: 50 ON

Low Mass Gate: 500.0

PSD Mirror Ratio:

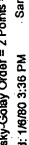
Negative Ions: OFF

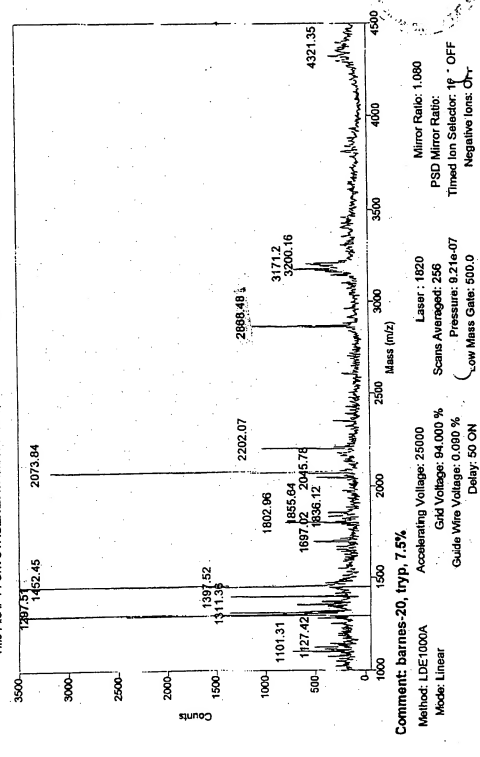


This File # 4 : C:\VOYAGERUDATAWAG100\DIGEST\SMOOTH.MS Original Filename: c:\voyager\data\mag100\digesf\bare001.ms

Collected: 1/6/80 3:36 PM

Savilsky-Golay Order = 2 Points = 19



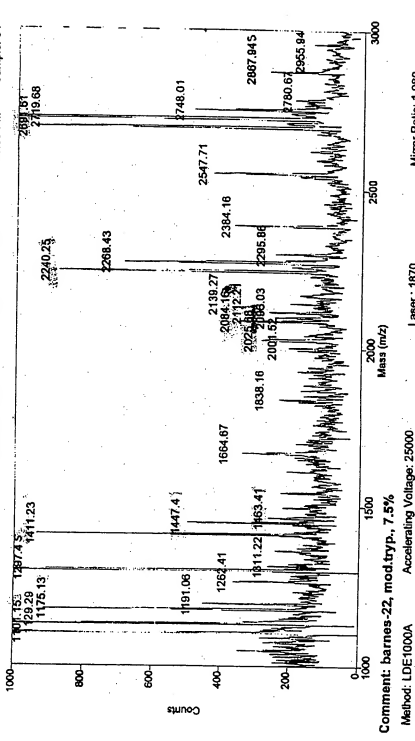


7



This File # 1: C:IVOYAGERIDATAIMAG200DIGESTISMOOTH.MS Original Filename: c:\voyager\data\rnag200\digest\barne003.ms

Sample: 54 Savitsky-Golay Order = 2 Points = 19 Collected: 2/16/80 3:35 PM



Timed Ion Selector: 16 1 OFF

Pressure: 3.43e-07

Laser: 1870

Scans Averaged: 231

Grid Voltage: 94.000 %

Mode: Linear

Guide Wire Voltage: 0.090 %

Delay: 50 ON

PSD Mirror Ratio:

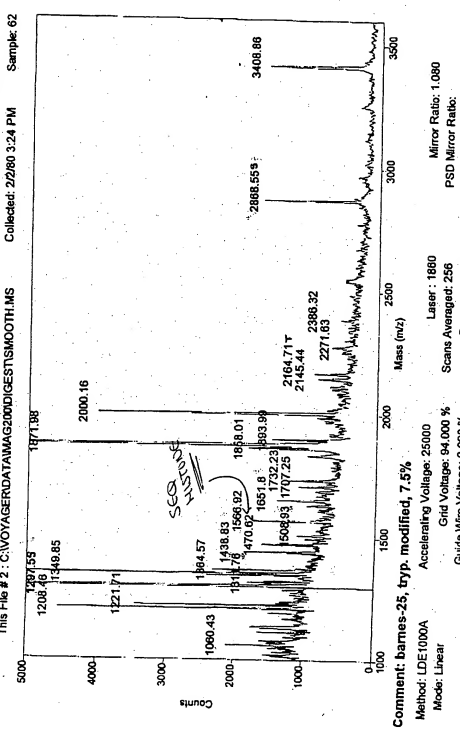
Negative lons: o

Mirror Ratio: 1.080





Savitsky-Golay Order = 2 Points = 19



Timed Ion Selector: 19 ' OFF

Pressure: 9.28e-07

Guide Wire Voltage: 0.090 %

Delay: 50 ON

LOW Mass Gate: 500.0

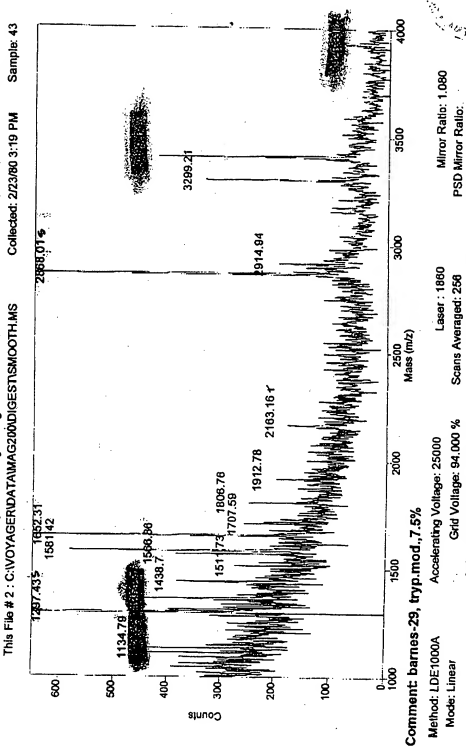
Negative lons: Orr

: ;

Columbia University /HHMI Protein Core

Original Filename: c:\voyager\data\mag200\digest\bame006.ms

Savitsky-Golay Order = 2 Points = 19 Collected: 2/23/80 3:19 PM



Timed Ion Selector: 16.1 OFF

Pressure: 3.82e-07

Guide Wire Voltage: 0.090 %

Delay: 50 ON

Low Mass Gate: 500.0

Negative lons: OFF

Figure 8

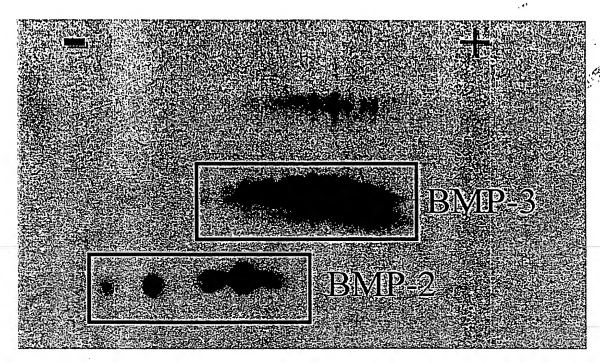


FIGURE 9A

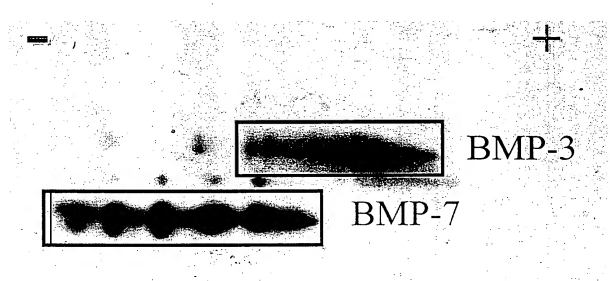


FIGURE 9B

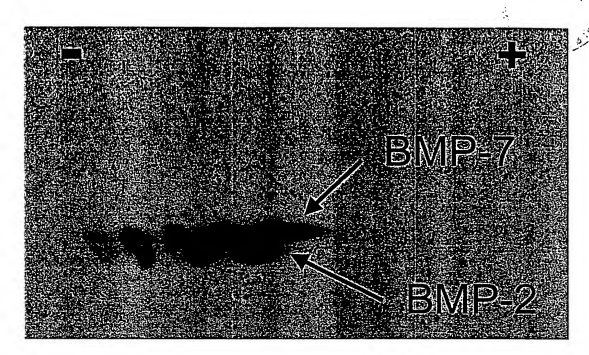


FIGURE 9C

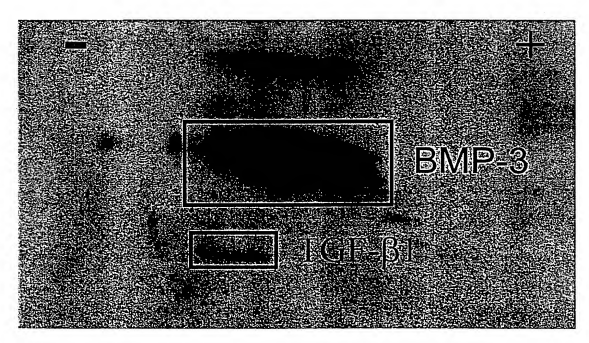
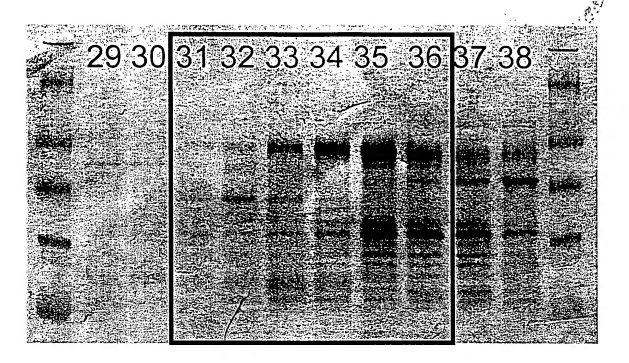
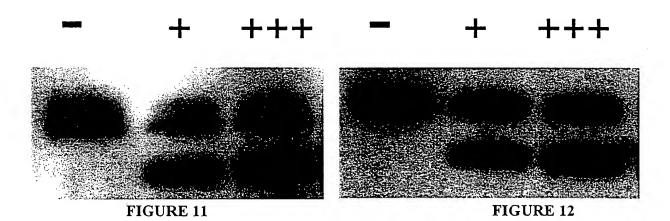
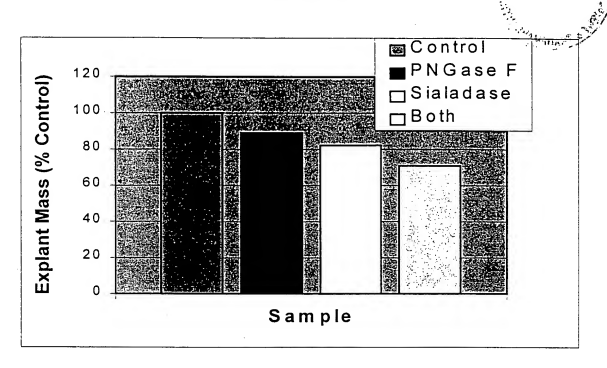


FIGURE 9D









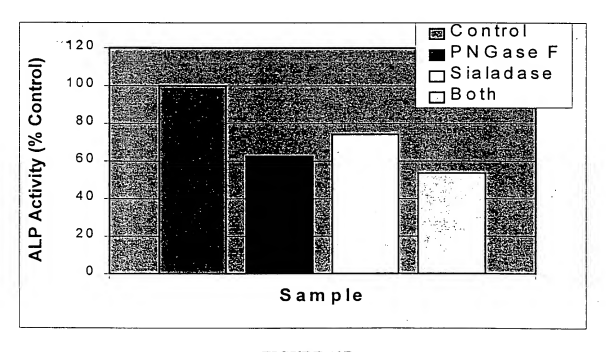


FIGURE 13B



Specificity	Antigen	Host Species	PC/MC	Source	Catalog No.
TGF-β1 (human)	Protein	Rabbit	Polyclonal	Promega	G1221
TGF-β2 (human)	Peptide	Rabbit	Polyclonal	Santa Cruz Biotechnology	sc-90
TGF-β3 (human)	Peptide	Rabbit	Polyclonal	Santa Cruz Biotechnology	sc-82
BMP-2 (human)	Protein	Rabbit	Polyclonal	Austral Biologics	PA-513-9
BMP-3 (human)	Peptide	Chicken	Polyclonal	Research Genetics	NA
BMP-4 (human)	Peptide	Goat	Polyclonal	Santa Cruz Biotechnology	so-6896
BMP-5 (human)	Peptide	Goat	Polyclonal	Santa Cruz Biotechnology	sc-7405
BMP-6 (human)	Peptide	Mouse	Monoclonal	Novocastra Laboratories	NCL-BMP6
BMP-7 (human)	Peptide	Rabbit	Polyclonal	Research Genetics	NA
FGF-1 (human)	Peptide	Goat	Polyclonal	Santa Cruz Biotechnology	sc-1884
osteonectin (bovine)	Protein	Mouse	Monoclonal	DSHB	AON-1
osteocalcin (bovine)	Protein	Rabbit	Polyclonal	Accurate Chemicals	A761/R1H
serum albumin (bovine)	Protein	Rabbit	Polyclonal	Chemicon International	AB870
transferrin (human)	Protein	Chicken	Polyclonal	Chemicon International	AB797
apo-A l lipoprotein (human)	Protein	Goat	Polyclonal	Chemicon International	AB740

Figure 15A: Identification of Proteins by Amino Acid Sequencing of Tryptic Fragments from 1D Gels

Band	Sample	Sequence Data	Best Database Match	Match	Identification	Species	Accession No.	AAs
 -								
2	fx 49 (1579)	fx 49 (1579) XLAAAGYDVEK	ALAAAGYDVEK	11/11	histone H1.c	human	human 87668 (NCBI)	65-75
က	fx 67 (1346)	fx 67 (1346) SLEKVCADLIR	SLEKVCADLIR	11/11	40s Ribosomal Protein S20	īat	R3RT20 (PIR)	31-41
4	fx 65 ()	fx 65 () (V)VCGMLGFPSEAPV	VVCGMLGFPGEKRV	11/14	LORP	mouse	AAC95338 (NCBI)	213-226
2	N terminal seq	N terminal seq STGVLLPLQNNELPG	STGVLLPLQNNELPG	15/15	BMP-3	human	human 4557371 (NCBI)	290-304
	fx 72 (3925)	fx 72 (3925) STGVLLPLQNNELPGAEYQY STGVLLPLQNNELPGAEYQY	STGVLLPLQNNELPGAEYQY	20/20	BMP-3	human	human 4557371 (NCBI)	290-309
:	fx 74 (3409) STGVLLPLQ	STGVLLPLQ	STGVLLPLQ	6/6	BMP-3	human	human 4557371 (NCBI)	290-298
9	fx 55 (1566)	fx 55 (1566) (S)QTLQFXE	SQTLQFDE	2//8	BMP-3	human	numan 4557371 (NCBI)	346-353
	fx 47	VYAF	no match		111			
	N terminal seq	N terminal seq HAGKYSREKNT(P)A(P)	HGGKYSREKNQPKP	11/14	α2-Macroglobulin Receptor Assoc. Pro.	human	human P30533 (Swiss-Prot)	31-46
	fx 57 (1438)	fx 57 (1438) SQTLQFDEQ	SQTLQFDEQ	6/6	BMP-3	human	4557371 (NCBI)	346-354
	fx 57 (1652) SLKPSNHA	SLKPSNHA	SLKPSNHA	8/8	BMP-3	human	4557371 (NCBI)	410-417
7	fx 51 (1093)	fx 51 (1093) AALRPLVKP	AALRPLVKP	6/6	60s Ribosomal Protein L32	monse	P17932 (Swiss-Prot)	1-9
i !	fx 37 (no MS)	fx 37 (no MS) A(H)I(Q)VERYV	AIVER	5/5	60s Ribosomal Protein L32	mouse	P17932 (Swiss-Prot)	109-113
	fx 37 (no MS)	fx 37 (no MS) A(H)I(Q)VERYV	HQSDRYV	2/2	60s Ribosomal Protein L32	mouse	P17932 (Swiss-Prot)	22-28
8	fx 78 ()	fx 78 () XALF(G)AQLGXALGPI	no match		777			
6	fx 56 (1567)	fx 56 (1567) SOTLOFDEOT	SQTLQFDEQT	10/10	BMP-3	human	P12645 (Swiss-Prot)	346-355



Figure 15B: Identification of Proteins by Amino Acid Sequencing of Tryptic Fragments from 1D Gels

Band	Band Sample	Sequence Data	Best Database Match	Match	Identification	Species	Species Accession No.	AAs
=	fx 55 (1311) SQTLXF	SQTLXF	SQTLQF	9/9	BMP-3	human	4557371 (NCBI)	346-351
:	fx 47 (1772)	VLATVTKPVGGDK	VLATVTKPVGGDK	13/13	60s Ribosomal Protein L6	human	human Q02878 (Swiss-Prot)	87-99
	fx 76 (1795)	xVFAL	VFAL	4/4	60s Ribosomal Protein L6	human	Q02878 (Swiss-Prot)	273-276
	fx 61 (1145)	fx 61 (1145) AVPQLQGYLR	AIPQLQGYLR	9/10	60s Ribosomal Protein L6	human	Q02878 (Swiss-Prot)	262-271
18								
22	fx 58 (1101) ALDAAYCFR		ALDAAYCFR	6/6	TGF-β2	human	human P08112 (Swiss-Prot)	303-311
:	fx 69 (no match)	CPYL	GYNANFCAGACPYL	14/14	TGF-ß2	human	numan P08112 (Swiss-Prot)	340-353
!	fx 66 (1411.71)		VNSQSLSPY	6/6	SPP24	bovine	Q27967 (Swiss-Prot)	42-50
25	fx 39 (1470)	KAAKPSV(P)	KAAKPSVP	8/8	Histone H1.x	human	JC4928 (PIR)	199-206
53								

fx≓fraction number (molecular weight of fragment, as measured by SDS-PACE)

Figure 16A: Identification of Proteins by Mass Spectrometry of Tryptic Fragments from ID Gels

0000	Maco Cross Brofile	00,000	Accession	Mass Spec	Mass Spec	Mass	944	O'COVORAGO	Comments
0	balla Mass spec riollie species	salpade	Number	Data	Database	Difference	n C	70 CO 4CI 89C	2
-	4 peaks match with	human	87668 (NCBI)	1172.97	1172.37	09:0	110-121	22	
	histone H1 c			1579.87	1579.71	0.16	62-29		15 MS peaks match with Band 2
				1708.47	1707.89	0.58	64-79		
				2011.58	2012.32	-0.74	35-54		
2	3 peaks match with	human	87668 (NCBI)	1579.76	1579.71	0.05	62-79*	16	identification of starred peptide
	histone H1.c			1708.02	1707.89	0.13	64-79		confirmed by sequence analysis
				2012.12	2012.32	-0.20			15 MS peaks match with Band 1
3	7 peaks match with	rat	R3RT20 (PIR)	1129.76	1129.40	0.36	50-59	62	
	nhosome S20			1156.21	1156.30	60.0-	76-83		
	20 000000000000000000000000000000000000			1334.46	1334.62	-0.16	56-66		
				1352.13	1351.58	0.55	88-99	-	
			-	1518.04	1517.77	0.27	9-21		
				1919.02	1919.19	-0.17	5-21		
				3404.02	3404.87	-0.85	88-119		
4	3 peaks match with	human	NP002309	1987.95	1988.27	-0.32	150-167	9	12 MS peaks match with Band 8
	I vevi Oxidase RP		(Swiss-Prof)	2410.35	2410.63	-0.28	648-669		
			/ / / / / / /	2610.57	2610.10	0.47	455-478		

Figure 16B: Identification of Proteins by Mass Spectrometry of Tryptic Fragments from ID Gels

Pac a	Mac Cnor Drofile	Sporioe	Accession	Mass Spec	Mass Spec	Mass	844	% Coverage	Comments
	Daild Mass Spec Floring Species	Species		Data	Database	Difference	ŝ	ognio o	
ည	9 peaks match with	human	4557371	1113.32	1113.31	0.01	361-368	48	% coverage calculation is relative
	RMP-3			1438.53	1438.58	•	346-357		to the mature BMP-3, 183 AAS
			(222)	1566.76	1566.76	00.00	345-357		(290-472)
_				1651.86	1651.91	-0.05	410-424		
				1794.09	1794.02	0.07	346-360		
			•	2268.46	2268.63	-0.17	374-392		
				2424.45	2424.81	-0.36	373-392		Identification of starred peptide
				3409.15	3407.77	1.38	290-318*		confirmed by sequence analysis
ဖ	3 peaks match with human	human	P30533	1002.24	1002.15	60.0	283-290	11	
	α2-Macroglobulin		(Swiss-Prot)	2362.58	2362.43	0.15	129-150		
	RAP			3048.51	3048.52	-0.01	257-282		
	2 peaks match with human	human	4557371	1566.93	1566.75	0.18	346-357	15	% coverage calculation is relative
	BMP-3		(NCBI)	1651.88	1651.91	-0.03	410-424	,	to the mature BMP-3, 183 AAS
									(290-472)

Figure 16C: Identification of Proteins by Mass Spectrometry of Tryptic Fragments from ID Gels

			Accession	Mass Spec	Mass Spec	Mass	900	% Coverage	Comments
Dand	band mass spec Prome species	salpade	Number	Data	Database	Difference	Š	76 00 to 18	
7	4 peaks match with	mouse	P17932	1033.25	1033.17	90.0	67-75	33	
	rihosome 1 32		(Swiss-Prof)	1093.31	1093.40	60'0-	1-10*		
			(2) (2)	1134.72	1134.28		65-74		
				1449.78	1449.66	0.12	19-29		
	5 peaks match with	human	4557371	1060.42	1060.20	0.22	102-111	21	% coverage catculation is relative to
-	BMD-3		(NCRI)	1113.39	1113.31	80.08	361-368		the mature BMP-3, 183 AAS (290-
	2-1810		(incori)	1360.26	1360.58	-0.32	190-200		472)
				1652.28	1651.91	0.37	410-424		
				1793.62	1794.02	-0.40	346-360		
ω	1 peak matches with	human	NP002309	2410.37	2410.63	-0.26	648-669	င	12 MS peaks match with Band 4
	Lysyl Oxidase RP		(Swiss-Prot)				:		
6	6 peaks match with	human	4557371	1113,14	1113.31	-0.17	361-368	36	% coverage calculation is relative to
٠	BMD-3		(NCRI)	1438.60	1438.58	0.02	346-357		the mature BMP-3, 183 AAS (290-
			(1001)	1566.77	1566.76	0.01	345-357		472)
				1651.91	1651.61	0.30	410-424		
				2901.67	2901.19		41-66		
	-			3408.94	3407.77	1.17	290-318		

Figure 16D: Identification of Proteins by Mass Spectrometry of Tryptic Fragments from ID Gels

Dang	Mac Caco Drofile	Casalas	Accession	Mass Spec	Mass Spec	Mass	9 4 6	% Coverage	Common
282	Dalld Mass Spec Fiolife	Species	Number	Data	Database	Difference	0	/ COVEIAGE	200
1	5 peaks match with	human	4557371	1113.23	1113.31	90.0-	361-368	48	% coverage calculation is relative to
	BMP-3		(NCRI)	1651.73	1651.91	-0.18	410-424		the mature BMP-3, 183 AAS (290-
	5		(1201)	1793.58	1794.02	-0.44	346-360		472)
				2424.24	2424.81	-0.57	373-392		
				3408.34	3407.77	0.57	290-318		
	5 peaks match with	human	Q02878	1140.38	1140.23	0.15	114-122	16	
	ribosome L6		(Swiss-Prot)	1526.88	1526.86	0.05	141-155		
		monse	P47911	1059.15	1059.12	0.03	10-20		
			(Swiss-Prof)	1145.36	1145.35	0.01	262-271		
			(20.000)	1386.74	1386.68	90.0	260-271		
18	4 peaks match with	human	P08112	1101.20	1101.26	90.0-	303-311	52	
	TGF-II2		(Swiss-Prof)	1175.26	1175.42	-0.16	400-409		
	<u> </u>		(20.1.2)	2240.37	2240.60	-0.23	312-328		
				2691.70	2691.91	-0.21	340-362		
	5 peaks match with	bovine	Q27967	1410.93	1411.60	-0.67	42-53	30	
	SPP24		(Swiss-Prot)	1447.59	1447.65	90.0-	113-124		
	5		(2011)	1540.64	1540.60	0.04	86-98		
				1869.10	1869.05	0.05	62-77		
				2268 47	2268 57	-0 10	33-53		

Figure 16E: Identification of Proteins by Mass Spectrometry of Tryptic Fragments from ID Gels

Commonte										*				% coverage calculation is relative to	the mature BMP-3,183 AAS (280-472)			
% Coverage	% coverage	63					11		14					31				
9 4	277	303-311	400-409	312-347	312-328	340-362	42-53	113-124	48-57	107-118	107-119	48-58	43-57	102-111	346-357	345-357	410-424	290-318
Mass	Difference	-0.11	-0.29	-0.26	-0.35	-0.30	-0.37	-0.25	90.0	-0.64	79.0-	-0.02	-0.74	0.23	0.25	0.16	-0.11	1.09
Mass Spec	Database	1101.26	1175.42	2084.42	2240.60	2691.91	1411.60	1447.65	1208.40	1222.35	1350.52	1364.59	1732.97	1060.20	1438.58	1566.76	1651.91	3407.77
Mass Spec	Data	1101.15	1175.13	2084.16	2240.25	2691.61	1411.23	1447.40	1208.46	1221.71	1349.85	1364.57	1732.23	1060.43	1438.83	1566.92	1651.80	3408.86
Accession	Number	P08112	(Swiss-Prof)	(2)			Q27967	(Swiss-Prot)	JC4928 (PIR)	,				4557371	(BCN)	(1201)		
Crocion	calpade	human					bovine		human					human				
Dand Mac Coop Brofile	Mass spec rione	5 peaks match with	TGF-R2	<u>!</u>			2 peaks match with	SPP24	5 peaks match with	histone H1 x				5 peaks match with	RMP.3) [
מים	משות	22							25									



Figure 16F: Identification of Proteins by Mass Spectrometry of Tryptic Fragments from ID Gels

Comments	% coverage calculation is relative to	the mature BMP-3,183 AAS (290-472)		
% Coverage	27			
AAs		346-357		290-318
Mass Difference	•	0.12		1.27
Mas	1113.31		1566.75	3407.77
Mass Spec Data	1113.22	1438.70	1566.86	3409.04
ion	4557371	(NCRI)	(1)	
Species	human			
Band Mass Spec Profile Species	29 4 peaks match with human	RMP-3)	
Band	59			

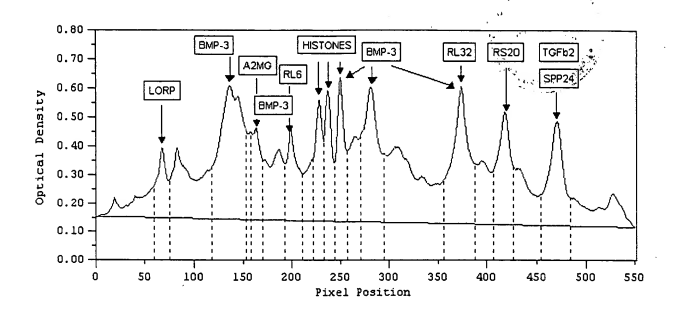


Figure 17A



Figure 17B

FIGURE 18: Quantitation of Identified BP proteins

Identified Protein	Percentage of Total Protein
LORP	2
BMP-3	11
BMP-3 and A2-MG	3
RL6 & BMP-3	4
Histone	3
Histone	3
Histone & BMP-3	4
BMP-3	8
RL32 & BMP-3	8
RS2D	5
SPP24 & TGF-β2	6
Total	58%

Figure 19A

Ideatification of Proteinsby Riass Spectrometry of Fragments from 2D Gels

Comments		·			populae malah confirmed by	sequence analysis											
% Coverage		9					ka		4			-	•				
AAS		472-487		364-382	488-504		241-253	645-668	105-116		58-70	ZI-33	301-314	36-134	774-295	239-261	131-154
	吾	Q. 13		0.51	NIA		.0.31	0.20	D.40		0.14	0.35	1.06	0.71	1.40	D. 50	D.43
	Darisbase	1837.14		1921.14	VN		1609.AB	2410.63	1606.60		1548.70	QU 0991	08.0481	1634.00	2351.50	2380.70	2727.50
NS Peaks	Dada	1857.01		1921.65	2679.51		1609.57	2410.89	1407.76	.:	1546.64	1681,16	18B1 A6	163671	0872587	2381.50	272(51
AEC. Na.		P05160 (Swiss- Prof)					NP002308 (SMCE-Prot)		P25075 (Swiss-	(ma.a	•						
Species		Human		· · .			Human	•	Bovine					- 14 - 14			
Mass Spec		2 peaks maich	Cospulation Factor XIIIb				Trypsin 2 peaks match		Lys-C & peaks metch	Win Centerior				٠. "			
Spel Digest		Lysc					Trypsin		Lysc								
25		-	. '				~		2								

Figure 19B

Identification of Proteinsby Mass Spectrometry of Fragments from 2D Gels

Commonte				peptide matches	confirmed by sequence analysis					•								•			i i	₩.		- 4		
72	2	Coverage					20			16		90					-				٠.					
AAc	}	٠	,				26-31	32-37	98-107	42-80	21-32	78-85			99-108	99-108	42-53	113-124	86-98	85-88	62-77	61-77	21-41	78-98	88-108	125-151
			Diff	N/N/N/N/N/N/N/N/N/N/N/N/N/N/N/N/N/N/N/		2.02	-0.34	-0.27	-0.17	-0.02	0.47	-0.09			-0.24	0.11	-0.07	-0.02	0.05	0.08	0.11	0.77	0.41	0.53	-0.51	0.79
			Database	N/A		4593.08	774.90	809.94	1175.43	1415.58	2187.51	1078.15			1101.31	1172.31	1411.60	1447.65	1540.52	1696.71	1869.05	2025.24	2272.56	2599.65	2693.81	2928.01
116	Ē	Peaks	Data	1461.58		4595.08	774.56	809.67	1175.28	1415.56	2187.98	1078.08			1101.07	1172.42	1411.53	1447.63	1540.57	1696.79	1889.18	2026.01	2272.97	2600.18	2693.30	2928.80
Ace No	ALL. (30.			P16836 (Swiss-	Prot)		P21214 (Swiss-			QZ7967 (Swiss-	Ê	QZ7967 (Swiss-	Prof)					•								
	Sheries			Rat			Bovine	••		Bovine		Bovine														
Г	ລ	Profile		2 peaks match	with Lysyl Oxidase		3 peaks match			2 peaks match	#7.1.0 Inu	13 peaks	match with	SPP24					•				·			
<u> </u>	Digest			Lys-C			Lys-C					Trypsin														
6	200			4		·	5			,		0														

Figure 19C Identification of Proteinsby Mass Spectrumetry of Wragments from 2D Gets

Comments																· · · · · · · · · · · · · · · · · · ·			100		
F	Coverage		42				10			37		.•								- :	
A			26-31	32-37	BB-107	1-25	42-60	:		340-355	10-18	285.206	249-260	103-114	103-115	84-49	30-49	177-197	200-223	70.90	199.223
		Diff	-0.34	-0.25	150-	1.44	0.20			0.25	0.0B	0.22	0.02	0.17	0.04	-0.16	-0.12	0.34	0.27	-0.49	-0.25
		Database	774.80	808.94	1175.43	3166.88	187.51		· .	917.14	984.15	1192.40	1380.65	1484.69	162D,82	1770.00	2238.55	39'5ZEZ	2561.04	2896.43	2948.35
SE SE	Pazks	Dada	774.55	809.63	1175.12	3168.10	2167.77			217.23	98(23	1192.62	1380.67	1484.60	1620.08	1770.64	2238.43	2125.99	16188	2697.84	2046.10
Ack. No.			P21214 (Swiss- Prol)				027867 (Swiss-	Pag.	•	P30072 (Swiss- Prof)						•	٠				
Species	٠.		Bowne				Bovine			Bovine										1.	
Muss Spec	Profile		4 peaks match with TGF-62				(beak	metches with SPP24		Trypsin 12 peaks match with abosome	3										
Spat Digest			Lysc						· .	Trypsin											
Spal			~							-											

Figure 19D

Identification of Proteinsby Mass Spectrometry of Bragments from 2D Gels

F	· · · · · · · · · · · · · · · · · · ·							11 Topsio			~	•••	opedul of				-	,			9 Typsin		redu nate	
							MM Honorous	<u> </u>			1	H1C Helvin	믓							with sidesokne	7 peaks match		Profile	
				·.				Human	,			:	Human			,					Mouse		Courselo	and in the
							1.09	P12750 (SHiss-				•	67656 (HCBI)							Pg.	P97351 (Swiss-			Acc No
	2591.60	2140.34	1587.75	1507.81	1354.03	1216.38		1188.40	2147.17	1707.65	1578.70		(327.75	1953,12	1719.91	1593.72	1516,69	1340.62	1218.20		20.020	Data	Peaks	SW
	2591.90	2140.58		(SDY.OB	1353.61	1216.39		1 (60.38		1707.89	1579.71		1377.58	1053.16	1720.00	1593.02	15(8,69	1346.49	1218.31	•	020,10	Dataluase		
	0 -0.10	-0.24	-0.27	0.12	. 0 <i>A</i> 2	0,00		0,10	T.		-0.01		0,18	1.	60.07	ð.10	0,00	E 5.0	-0.02		-0.05	DAT		
1	7-9	221-23	_	108-21	230-24			230-23	7		65-78		SP-VC	65-81	199-212	94-108	174-186	151-161	152-181		19-26			A
	8	Τæ	1.0	T.E.				- 23					A.G		• .						29		Coverage	*
																								Comments